Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
Li	2830	544/114, 544/238, 544/239, 514/236.5, 514/252.02, 514/252. 03, 514/252.04, 514/252.05, 514/252.06	US-PGPUB; USPAT	OR	OFF	2005/10/19 14:55
L2	1831	CDK2	US-PGPUB; USPAT	OR	OFF	2005/10/19 14:53
L3	39	I1 and I2	US-PGPUB; USPAT	OR	OFF	2005/10/19 14:53
L4	2130	544/114, 544/238, 544/239, 514/236.5, 514/252.02, 514/252. 03, 514/252.04, 514/252.05, 514/252.06	USPAT	OR	OFF	2005/10/19 14:55

chain bonds :

1-8 4-7 5-18 6-19 9-10 10-11 12-13 13-14

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6

exact/norm bonds :

1-2 1-6 1-8 2-3 3-4 4-5 4-7 5-6 5-18 9-10 10-11 12-13 13-14

exact bonds :

6-19

G2:[*1],[*2]

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:Atom 9:CLASS 10:CLASS

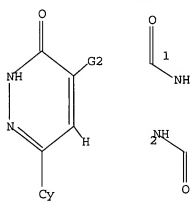
11:CLASS 12:CLASS 13:CLASS 14:CLASS 18:CLASS 19:CLASS

L1 STRUCTURE UPLOADED

=> d 11

L1 HAS NO ANSWERS

L1 STR



G1

G2 [@1],[@2]

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 08:42:39 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 59 TO ITERATE

100.0% PROCESSED 59 IT

59 ITERATIONS

9 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 720 TO 1640 PROJECTED ANSWERS: 9 TO 360

L2 9 SEA SSS SAM L1

<10/19/2005> Habte

=> s 11 sss full FULL SEARCH INITIATED 08:42:46 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 1173 TO ITERATE

100.0% PROCESSED 1173 ITERATIONS 103 ANSWERS

SEARCH TIME: 00.00.01

L3 103 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY SESSION

FULL ESTIMATED COST 161.33 161.54

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L4 24 L3

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L4 ANSWER 1 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 2005:1004734 CAPLUS DOCUMENT NUMBER: 143:306326

143:306326 Production of 4-benzimidazol-2-yl-pyridazin-3-one derivatives and use thereof in medicaments Schoenafinger, Karl: Hoelder, Swen; Will, David William; Hatter, Hams; Rueller, Guenther; Bossart, TITLE:

INVENTOR (S):

Walting Hatter, Hansy Hueller, Quenthers so Hartin Aventis Pharma Deutschland G.m.b.H., Germany PCT Int. Appl., 126 pp. CODEN: PIXXD2 Patent PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PRIORITY APPLN. INFO .:

PATENT NO.																				
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SY, TJ, TH, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZH, Z RY: BY, CH, CH, KE, LS, HY, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZY, AM, AZ, BY, KG, KZ, HD, RU, TJ, TH, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HJ, IZ, IS, IT, LT, LU, HC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CH, GA, GN, GQ, GY, HL, MR, NE, SN, TD, TG DE 102004010194 A1 20051013 DE 2004-102004010194 20040302				LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MV,	MX,	MZ,	NA,	NI,	
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				MR,	NE,	SN,	TD,	TG												
ORITY APPLN. INFO.: DE 2004-102004010194A 20040302	I	Œ	1020	0401	0194		A1		2005	1013		DE 2	2004-	1020	0401	0194	2	0040	302	
	ORI	T	APP	LN.	INFO	.:						DE 2	004-	1020	0401	0194	A 2	0040	302	

L4 ANSWER 1 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) deprotection of V (Y2 = protecting group). Thus, 4-(6-trifluoromethyl-lH-benzimidazol-2-yl)-6-(pyridin-4-yl)-2H-pyridazin-3-one [I] A = B = E = CH, D = C-CF3, Rl = 4-pyridinyl, R2 = H] was prepd. from 3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazin-4-carboxylic acid via chlorination with SOCl2 in (ReOCH2),2 followed by amination with 4-(trifluoromethyl)benzene-1,2-diamine in (ReOCH2) 2 contg. Et3N and cyclocondensation of the amide in AcOH. Said compds. are kinase inhibitors, particularly inhibitors of kinase GSK-39 (glycogen synthase kinase-39). The enzyme inhibitory activity of I (A = B = E = CH, D = C-CF3, Rl = 4-pyridinyl, R2 = H] was detd. [ICS0 = 16 nM].

IT 86464-06-2P
RL: RCT (Reactant): SPN (Synthetic preparation): PREP (Preparation): RACT (Reactant or reagent)
(preparation and cyclocondensation of: preparation of 4-benzimidazol-2-ylpyridazin-3-one derivs. with GSK-3β inhibitory activity)
RN 86464-06-2 CAPIUS

<10/19/2005>

Habte

The invention relates to compds. I [A = CR3, N, B = CR4, N, D = CR5, N, E = CR6, N, R1 = halogen, un-, monosubstituted Cl-10-alkyl, heterocyclyl, aryl, heterocaryl (optionally substituted with halogen, CN, NO2, OR7, COR7, CO2A7, OC(10)R7, NRTR8, NRCOR7, CONR7,88, NHCSR7, CSR7, SOR7, SOZA7, NHSOZR7, SOZAR7R8, OSCAR7R8, NRCSR7, SOZAR7R8, OSCAR7, SOZAR7, SOZ-OR7, aryl, heterocaryl, heterocycle, CR3, CCCT3), R2 = H, Cl-10-alkyl, R3, R4, R5, R6 = H, halogen, CN, NO2, CH2R8, CH2NH2, CH2NH(Cl-6-alkyl), CH2N(Cl-6-alkyl), OR8, COR8, COZR8, OC(10)R8, NR7R8, NRCOR8, CORR8, CSNR7R8, SSR8, SOR8, SOZR8, NHSOZR8, SOZNR7R8, OSCAR7R8, NRCSR8, CSNR7R8, SSR8, SOR8, SOZR8, NHSOZR8, SOZNR7R8, OSCAR7R8, NRCOR8, CSNR7R8, SSR8, SOR8, SOZR8, NHSOZR8, SOZNR7R8, OSCAR7R8, NRCOR8, CSNR7R8, NRCOR8, CSNR7R8, NRCOR8, CSNR7R8, SSR8, SOR8, SOZR8, NHSOZR8, SOZNR7R8, OSCAR7R, R8 = H, un-monosubstituted cl-10-alkyl, C2-10-alkynyl, heterocyclyl, aryl, heterocryl, aryl = 5- to 10-membered aromatic mono- or bicyclic ring, heteroaryl = 5- to 10-membered aromatic mono- or bicyclic ring with one or more heteroaroms - N, O, SI heterocycle = 5- to 10-membered non-aromatic mono- or bicyclic ring with one or more heteroaroms - N, O, SI with the proviso that up to three of A, B, D, E can equal N simultaneously; etc.] in addition to the physiol. compatible salts thereof, methods for the production of said compds. and the use thereof as medicaments. The procedure for the paration of 1 comprises: reaction of pyridazinone II [Y = H, leaving group] with

aration
of I comprises: reaction of pyridazinone II [Y = H, leaving group] with
diamine III whereby cyclization takes place (a) in the presence of an acid
or H2O removing medium when Y = leaving group or (b) through exidation,
or ally

or H20 removing medium when I = leaving gloup of th, chicago cially in the presence of O2, when Y = H. Alternatively, I can be prepared from pyridazin-3-one IV (YI = halogen, B(OH)2, Sn(Cl-10-alkyl)3; Y2 = H, protecting group) via palladium-catalyzed coupling with R12 [Z = halogen, B(OH)2, B(Cl-10-alkyl)2, Sn(Cl-10-alkyl)3, Zn(Cl-10-alkyl)] followed by

ANSWER 1 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

REFERENCE COUNT: THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSVER 2 OF 24
ACCESSION NUMBER:
DOCUMENT NUMBER:
11TILE:
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:

CAPIUS COPYRIGHT 2005 ACS on STN
2005:1002888 CAPIUS
103:286437
Preparation of 4-benzimidezol-2-yl-pyridazin-3-ones as cyclin dependent kinase 2 inhibitors
Aventis Pharma S. A., Fr.
COIDEN: GMOXEX
DOCUMENT TYPE:

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATE	ent i	NO.			KIN	D	DATE			APPL					D.	ATE		
						-									•			
DE 1	1020	0401	0207		A1		2005	0915		DE 2	004-	1020	0401	0207	2	0040	302	
WO 2	2005	0852	31		Al		2005	0915	,	WO 2	005-	EP25	69		2	050	218	
	v:	AE,	λG,	AL,	AH,	AT,	AU,	λZ,	BA,	BB,	BG,	BR,	BW.	BY,	BZ,	CA,	CH,	
		CN,	co.	CR,	CU,	CZ,	DE,	DK.	DM,	DZ,	EC,	EE,	EG.	ES,	FI.	GB,	GD,	
							ID.											
		LK.	LR.	LS.	LT.	LU.	LV.	MA.	MD.	MG.	MK.	MN,	MV.	HX,	MZ.	NA.	NI.	
		NO.	NZ.	OM.	PG.	PH.	PL.	PT.	RO.	RU.	sc,	SD.	SE,	SG.	SK.	SL.	SM.	
		SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	υs,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RV:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
		AΖ,	BY,	KG,	KZ,	MD,	RU,	IJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IS,	IT,	LT,	LU,	MC,	NL,	PL,	PT,	
		RO,	SE,	SI,	SK,	TR,	BF,	BJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	
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PRIORITY GI	APP	LN.	INFO	.:						DE 2	004-	1020	0401	0207	A 2	0040	302	

Title compds. I (A = CR3, N; B = CR4, N; D = CR5, N; E = CR8, N; R3, R4,

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
11TLE:
2004:411321 CAPLUS
140:423683
Preparation of pyridazinones as protein Tau
phosphorylation inhibitors, their drugs and
pharmaceutical compositions containing them for
treatment, in particular, of central and peripheral
nervous system diseases
Lesuisse, Dominique, Halley, Franck, Baudoin, Bernard,
Rooney, Thomas, Hoelder, Swen, Naumann, Thorsten,
Tiraboochi, Gilles
SOURCE:
DOCUMENT TYPE:

DOCUMENT TYPE:

CAPLUS COPYRIGHT 2005 ACS on STN
2004:411321 CAPLUS
140:423683
Preparation of pyridazinones as protein Tau
phosphorylation inhibitors, their drugs and
pharmaceutical compositions containing them for
treatment, in particular, of central and peripheral
nervous system diseases
Lesuisse, Dominique, Halley, Franck, Baudoin, Bernard,
Rooney, Thomas, Hoelder, Swen, Naumann, Thorsten,
Tiraboochi, Gilles
COEN: FRXXBL
Patent

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DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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PR	2847	253			A1		2004	0521		FR 2	002-	1444	3		20	0021	119	
CA	2506	022			A1 AA		2004	0603		CA 2	003-	2506	022		20	0031	119	
WO	2004	0461	30		A1		2004											
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		CN.	co.	CR.	CU.	CZ.	DE.	DK.	DM.	DZ.	EC.	EE.	EG.	ES.	FI.	GB.	GD.	
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										TO 2	002	2012:	250		7 20	1030	10/	
										= U 2	003-	GF 12:	990	,	- 20	1031	119	

OTHER SOURCE(S): MARPAT 140:423683 ANSWER 2 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
R5 = H, halo, CN, etc.; R1 = halo, alkyl; R2 = H, alkyl; R8 = H, alkyl,
alkenyl, etc.] and their pharmaceutically acceptable salts and
formulations were prepd. For example, sapon. of Me ester II (X = CMe)
afforded claimed carboxylic acid II (X = CM). In cyclin dependent kinase
2 inhibition assays, 3-examples of compds. I exhibited IC50 values ranging
from 0.026-0.214 µH.
684464-01-7P 864464-06-2P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(preparation of benzimidazolylpyridazinones as cyclin dependent kinase 2
inhibitors)
864464-01-7 CAPLUS
INDEX NAME NOT YET ASSIGNED

864464-06-2 CAPLUS INDEX NAME NOT YET ASSIGNED

ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN L4 (Continued)

Title compds. I [wherein A = CONHR, or NHCOR; R = (un)substituted heteroaryl/aryl/alkyl, hetero/aryl, fused hetero/aryl with cycloalkyl, etc.; Ar = (un)substituted aryl, Ph, pyridinyl; and their racemates, enantiomers, disstereomers, mixts., tautomers and pharmaceutically acceptable salts] were prepared as protein Tau phosphorylation inhibitors. Three standard pharmaceutical compns. are given. For example, II was ared

Three standard pharmaceutical compns. are given. For example, II was prepared by acylation of 3-0xo-6-phenyl-2,3-dihydropyridazine-4-carboxylic acid with 2,4-dichlorobenzylamine. Selected invention compds. I inhibited phosphorylation of protein Tau with an IC50 < 100 µM. Thus, I and their pharmaceutical compns. are useful as kinase inhibitors and for treatment, in particular, of central and peripheral nervous system diseases (no data).

IT 691868-73-69. N-1(2,4-Dichlorobenzyl)-3-oxo-6-[4-(benzyloxy)phenyl]-2,3-dihydropyridazine-4-carboxamide 691868-77-69. N-Benzyl-3-oxo-6-[4-(bydroxy)phenyl]-2,3-dihydropyridazine-4-carboxamide RL: RCT (Reactant): FSN (Synthetic preparation): PREP (Preparation): RACT (Reactant or reagent)

(Intermediate: preparation of pyridazinones as protein Tau phosphorylation

inhibitors for treating central and peripheral nervous system diseases)

NO 691848-75-6 CAPUS

CN 4-Pyridazinecarboxamide, N-[(2,4-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-6-(4-(phenylmethoxy)phenyl)- (9CI) (CA INDEX NAME)

11

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

691848-77-8 CAPLUS 4-Pyridazinecarboxami (phenylmathyl)- (9CI) -amide, 2,3-dihydro-6-(4-hydroxyphenyl)-3-охо-N-CI) (CA INDEX NAME)

691846-21-2P, N-(2,4-Dichlorobenzyl)-3-oxo-6-phen-4-yl-2,3-dihydropyridazine-4-carboxamide 691848-24-5P,
N-(2,4-Dichlorobenzyl)-3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide 691848-21-3P,
N-(2,4-Dichlorobenzyl)-3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide 691848-31-4P,
N-(4-Chlorobenzyl)-3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide 691848-33-6P,
N-(2-(2)-4-Dichlorophenyl) sthyl]-3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide 691848-36-9P,
N-(2,4-Dichlorophenyl) sthyl]-3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide 691848-36-9P,
N-(2,4-Dichlorophenyl)-3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide 691848-31-6P,
3-Oxo-6-(pyridin-4-yl)-N-(3-(trifluoromethyl) benzyll-2,3-dihydropyridazine-4-carboxamide 691848-43-6P,
3-Oxo-6-(pyridin-4-yl)-N-(3-bihydropyridazine-4-carboxamide
691848-31-2P,
N-(3,5-bihclorobenzyll-3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide
691848-31-8P,
3-Oxo-6-(pyridin-4-yl)-N-(n-butyl)-2,3-dihydropyridazine-4-carboxamide
691848-31-8P,
3-([3-oxo-6-(pyridin-4-yl)-N-(pyridin-3-yl)-2,3-dihydropyridazin-4-carboxamide
691848-31-8P,
3-([3-oxo-6-(pyridin-4-yl)-N-(pyridin-4-yl)-2,3-dihydropyridazin-4-carboxamide
691848-31-8P,
3-([3-oxo-6-(pyridin-4-yl)-N-(pyridin-4-yl)-2,3-dihydropyridazin-4-carboxamide
691848-31-8P,
3-([3-oxo-6-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-N-(pyridin-4-yl)-2,3-dihydropyridazine-4-carboxamide
691848-31-8P,
3-([0-oxo-6-(pyridin-4-yl)-N-(pyridin-4-yl)-3-(pyridin-4-yl)-N-(pyridin-4-y

ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
691849-29-3P, 3-Oxo-6-[4-(hydroxy)pyridin-3-yl]-N-(pyridin-2-ylmethyl)-2,3-dihydropyridazin-e-4-carboxanide 691849-30-6P,
N-(3,4-Dichlorobenzyl)-3-oxo-6-[4-(hydroxy)pyridin-3-yl]-2,3dihydropyridazin-e-4-carboxanide 691849-31-7P,
N-(4-(Morpholin-4-yl)benzyl]-3-oxo-6-[4-(hydroxy)pyridin-3-yl]-2,3dihydropyridazin-e-4-carboxanide 691849-32-6P,
N-(4-Hydroxybenzyl)-3-oxo-6-[4-(hydroxy)pyridin-3-yl]-2,3dihydropyridazin-e-4-carboxanide 691849-33-9P,
2-(2,4-Dichlorophenyl)-N-[3-oxo-6-(pyridin-4-yl)-2,3-dihydropyridazin-4yl]-3-dihydropyridazin-e-4-carboxanide
RL: PAC (Pharmacological activity) SFN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)

(Uses)
(protein Tau phosphorylation inhibitor; prepn. of pyridazinones as protein Tau phosphorylation inhibitors for treating central and peripheral nervous system diseases)
691848-21-2 CAPLUS
4-Pyridazinecarboxamide, N-[(2,4-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-6-phenyl- (9CI) (CA INDEX NAME)

691848-24-5 CAPLUS 4-Pyridazinecarboxamide, N-[(2,4-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

691848-28-9 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-N-(phenylmethyl)-6-(4-pyridayi)-(9CI) (CA INDEX NAME)

ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) carboxamide 691848-59-6F, N-[4-(Morpholin-4-yl)benzyl]-3-oxo-6-(pyridin-4-yl)-2, 3-dihydropyridazine-4-carboxamide 691848-67-6F, N-[4-Nydroxybenzyl]-3-oxo-6 (pyridin-4-yl)-2, 3-dihydropyridazine-4-carboxamide 691848-19-0F, N-[2,4-Dichlorobenzyl]-3-oxo-6-[4-(hydroxyl)penyl]-2, 3-dihydropyridazine-4-carboxamide 691848-19-0F, N-[2,4-Dichlorobenzyl]-3-oxo-6-(pyridin-2-yl)-2,3-dihydropyridazine-4-carboxamide 691848-19-0F, N-[2,4-Dichlorobenzyl]-3-oxo-6-(pyridin-2-yl)-2,3-dihydropyridazine-4-carboxamide 691849-9-4F, N-[2,4-Dichlorobenzyl]-3-oxo-6-(pyridin-3-yl)-2,3-dihydropyridazine-4-carboxamide 691849-03-3F, N-[4-Chlorobenzyl]-3-oxo-6-[4-(hydroxy)phenyl]-2,3-dihydropyridazine-4-carboxamide 691849-04-4F, N-[2-Chlorobenzyl]-3-oxo-6-[4-(hydroxy)phenyl]-2,3-dihydropyridazine-4-carboxamide 691849-05-5F, N-[2-(2,4-Dichlorophenyl)-3-oxo-6-[4-(hydroxy)phenyl]-2,3-dihydropyridazine-4-carboxamide 691849-07-7F, 3-oxo-6-[4-(hydroxy)phenyl]-2,3-dihydropyridazine-4-carboxamide 691849-07-7F, 3-oxo-6-[4-(hydroxy)phenyl]-N-[pyridazine-4-carboxamide 691849-07-7F, 3-oxo-6-[4-(hydroxy)phenyl]-3-(3-0xo-6-[4-(hydroxy)phenyl]-3-(3-1)-(4-(hydroxy)phenyl)-3-(3-1)-(4-(hydroxy)phenyl)-3-(3-1)-(4-(hydroxy)phenyl)-3-(3-1)-(4-(hydroxy)phenyl)-3-(3-1)-(4-(hydroxy)phenyl)-3-(3-1)-(4-(hydroxy)phenyl)-3-(3-1)-(4-(hydroxy)phenyl)-3-0xo-6-(4-(hydroxy)phenyl)-3

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

691848-31-4 CAPLUS 4-Pyridazinecarboxamide, N-{(4-chlorophenyl)methyl}-2,3-dihydro-3-oxo-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

691848-33-6 CAPLUS 4-Pyridazinecarboxamide, N-[(2-chlorophenyl)methyl]-2,3-dihydro-3-oxo-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

691848-36-9 CAPLUS 4-Pyridazinecarboxamide, N-{2-(2,4-dichlorophenyl)ethyl]-2,3-dihydro-3-oxo-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

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L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

691840-39-1 CAPUS 4-Pyridarinecarboxamide, N-(2,4-dichlorophenyl)-2,3-dihydro-3-oxo-6-(4-pyridinyl)- (9C1) (CA INDEX NAME)

691848-41-6 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-N-(4-pyridinylmethyl)- (9CI) (CA INDEX NAME)

691848-43-8 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-N-[{3-(trifluoromethyl)phenyl]methyl]- (9CI) (CA INDEX NAME)

ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

691848-51-8 CAPLUS \$\textit{\beta}\-Alanine, N-\[(2,3-\text{dibydro-3-oxo-6-(4-pyridinyl)-4-pyridazinyl\]carbonyl\]-, ethyl ester (9CI) (CA INDEX NAME)

691848-53-0 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-N-(3-pyridinylnethyl)- (9CI) (CA INDEX NAME)

691848-55-2 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-N-(2-pyridinyl)-N-(CA INDEX NAME)

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L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

691848-45-0 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-N-[[4-trifluoromethyl]phenyl]methyl]- (9CI) (CA INDEX NAME)

691848-47-2 CAPLUS
4-Pyridazinecarboxamide, N-[(3,5-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

691848-49-4 CAPLUS 4-Pyridazinecarboxamide, N-butyl-2,3-dibydro-3-oxo-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) 691848-57-4 CAPLUS 4-Pyridazinearboxamide, N-[(3,4-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

691848-59-6 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-N-[[4-(4-morpholiny1)phenyl]methyl]-3oxo-6-(4-pyridiny1)- (9CI) (CA INDEX NAME)

691848-67-6 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-N-[(4-hydroxyphenyl)methyl]-3-oxo-6-(4-pyridinyl)- (9Cl) (CA INDEX NAME)

691848-79-0 CAPLUS
4-Pyridazinecarboxamide, N-[{2,4-dichlorophenyl}methyl}-2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

691848-81-4 CAPLUS
4-Pyridazinecarboxamide, N-[(2,4-dichlorophenyl)methyl]-2,3-dihydro-6-[4-hydroxy-3-(phenylmethyl)phenyl]-3-oxo- (9CI) (CA INDEX NAME)

691848-89-2 CAPLUS
4-Pyridazinecarboxamide, N-[(2,4-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-6-(2-pyridinyl)- (9CI) (CA INDEX NAME)

- 691848-99-4 CAPLUS
 4-Pyridazinecarboxamide, N-[(2,4-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-
- ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) 691849-05-5 CAPLUS 4-Pyridazinearaboxamide, N-[2-(2,4-dichlorophenyl)ethyl]-2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo- (9CI) (CA INDEX NAME)

691849-06-6 CAPLUS 4-Pyridazi necarboxamide, N-(2,4-dichlorophenyl)-2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo- (9CI) (CA INDEX NAME)

691849-07-7 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo-N-(4-pyridinylmethyl)- (3C1) (CA INDEX NAME)

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ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN 6-(3-pyridinyl)- (9CI) (CA INDEX NAME) (Continued)

691849-03-3 CAPLUS 4-Pyridazinecarboxamide, N-{(4-chlorophenyl)methyl}-2,3-dihydro-6-{4-hydroxyphenyl)-3-oxo- (9Cl) (CA INDEX NAME)

691849-04-4 CAPLUS
4-Pyridazinecarboxamide, N-[(2-chlorophenyl)methyl]-2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo- (9CI) (CA INDEX NAME)

- L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
- 691849-08-8 CAPLUS
 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo-N-[{3-(trifluoromethyl)phenyl]methyl}- (9CI) (CA INDEX NAME)

691849-09-9 CAPLUS
4-Pyridazi necarboxamide, 2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo-N-[[4-(trifluoromethyl)phenyl]methyl]- (9CI) (CA INDEX NAME)

691849-10-2 CAPLUS
4-Pyridazinecarboxamide, N-[(3,5-dichlorophenyl)methyl]-2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-11-3 CAPLUS
CN 4-Pyridazinecarboxamide, N-butyl-2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo(9C1) (CA INDEX NAME)

RN 691849-12-4 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo-N-(3-pyridinylmethyl)- (9CI) (CA INDEX NAME)

RN 691849-13-5 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo-N-(2-pyridinylmethyl)- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-17-9 CAPLUS
CN 4-Pyridazinecarboxamide, N-[(2,4-dichlorophenyl)methyl]-2,3-dihydro-6-(4-hydroxy-3-pyridnyl)-3-oxo- (9CI) (CA INDEX NAME)

RN 691849-18-0 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxy-3-pyridiny1)-3-oxo-N-(phenylmethy1)- (9C1) (CA INDEX NAME)

RN 691849-19-1 CAPLUS
CN 4-Pyridazinecarboxamide, N-[(4-chlorophenyl)methyl]-2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-14-6 CAPLUS CN 4-Pyridazinecarboxamide, N-{(3,4-dichlorophenyl)methyl}-2,3-dihydro-6-(4-hydroxyphenyl)-3-oxo- (9CI) (CA INDEX NAME)

RN 691849-15-7 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dibydro-6-(4-hydroxyphenyl)-N-[(4-(4-morpholinyl)phenyl)methyl]-3-oxo- (9CI) (CA INDEX NAME)

RN 691849-16-8 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxyphenyl)-N-[(4-hydroxyphenyl)methyl)-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-20-4 CAPLUS
CN 4-Pyridazinecarboxamide, N-[(2-chlorophenyl)methyl]-2,3-dihydro-6-{4-hydroxy-3-pyridinyl)-3-oxo- (9CI) (CA INDEX NAME)

RN 691849-21-5 CAPLUS
CN 4-Pyridazinecarboxamide, N-[2-(2,4-dichlorophenyl)ethyl]-2,3-dihydro-6-(4-hydroxy-3-pyridainyl)-3-oxo-(9CI) (CA INDEX NAME)

$$\begin{array}{c} C1 \\ CH_2-CH_2-NH-C \\ \hline \\ N \\ N \\ \end{array}$$

RN 691849-22-6 CAPLUS
CN 4-Pyridazinecarboxamide, N-(2,4-dichlorophenyl)-2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo-(9CI) (CA INDEX NAME)

<10/19/2005>

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Page 11

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-23-7 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo-N-(4-pyridinylaethyl)- (9CI) (CA INDEX NAME)

RN 691849-24-8 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo-N-[[3-(trifluoromethyl)phenyl]methyl]- (9CI) (CA INDEX NAME)

RN 691849-25-9 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo-N-[[4-(trifluoromethyl)phenyl]methyl]- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-29-3 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo-N-(2-pyridinylmethyl)-(9CI) (CA INDEX NAME)

RN 691849-30-6 CAPLUS
CN 4-Pyridazinearboxamide, N-[(3,4-dichlorophenyl)methyl]-2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo-(9CI) (CA INDEX NAME)

RN 691849-31-7 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-N-[[4-(4-morpholinyl)phenyl]methyl]-3-oxo-(SCI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-26-0 CAPLUS
CN 4-Pyridazinecarboxamide, N-[(3,5-dichloropheny1)methy1]-2,3-dihydro-6-(4-hydroxy7-pyridiny1)-3-oxo- (9CI) (CA INDEX NAME)

RN 691849-27-1 CAPLUS
CN 4-Pyridazinecarboxamide, N-butyl-2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3oxo (9C1) (CA INDEX NAME)

RN 691849-28-2 CAPLUS
CN 4-Pyridazinecarboxanide, 2,3-dihydro-6-(4-hydroxy-3-pyridinyl)-3-oxo-N-(3-pyridinylnethyl)- (9Cl) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 691849-32-8 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-N-{(4-hydroxyphenyl)methyl]-6-(4-hydroxy-3-pyridinyl)-3-oxo- (9CI) (CA INDEX NAME)

RN 691849-33-9 CAPLUS CN Benzeneacetamide, 2,4-dichloro-N-[2,3-dihydro-3-oxo-6-(4-pyridinyl)-4pyridazinyl)- (9C1) (CA INDEX NAME)

RN 691849-34-0 CAPLUS
CN 4-Pyridazinecarboxamide, N-{(2,4-dichlorophenyl)methyl]-2,3-dihydro-3-oxo-6-(4-pyrimidinyl)- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSVER 4 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 2004:205958 CAPLUS DOCUMENT NUMBER: 142:93705 DOCUMENT NUMBER: TITLE: Product class 8: pyridazines Haider, N.: Holzer, W. AUTHOR (5): Halder, N., Holzer, W. Germany Science of Synthesis (2004), 16, 125-249 CODEN: SSCYJ9 Georg Thieme Verlag Journal, General Review CORPORATE SOURCE: SOURCE: UAGE: Journal, General Review
English
A review. Methods of preparing pyridazines are reviewed including
cyclization, ring transformation, aromatization, and substituent
modification.
87769-56-0 PUBLI SHER: DOCUMENT TYPE: LANGUAGE: AB A review. RI: RCT (Reactant): RACT (Reactant or reagent)
(review of preparation of pyridazines via cyclization, ring
transformation,
aromatization, and substituent modification)
RN 87769-56-0 CAPIUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-phenyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

INVENTOR (5):

THERE ARE 720 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE 720

L4 ANSWER 5 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 2003:293627 CAPLUS DOCUMENT NUMBER: 139:94783 139:94783
S-Aryl-pyrazolo[3,4-b]pyridazines: potent inhibitors of glycogen synthase kinase-3 (GSK-3)
Witherington, Jason: Bordas, Vincent: Haigh, David; Hickey, Deirdre M. B.: Ife, Robert J., Rawlings, Anthony D.; Slingsby, Brian P.; Smith, David G.; Ward, Robert W. AUTHOR (S):

SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

OTHER SOURCE(S):

Hickey, Deirdre M. B.: Ife, Robert J.: Rawlings,
Anthony D.: Slingsby, Brian P.: Smith, David G.;
Robert W.

PORATE SOURCE: Neurology Centre of Excellence for Drug Discovery,
Department of Medicinal Chemistry, GlaxoSmithXline
Research Limited, Harlow, CM19 SAW, UX
13(9), 1581-1594
CODEN: EMCLES: ISSN: 0960-894X
LISHER: Elsevier Science B.V.
MENT TYPE: Journal
IUAGE: English
Introduction of a nitrogen atom into the 6-position of a series of
Pyrazologi, 4-bjpyridines led to a dramatic improvement in the potency of
GSK-3 inhibition. Rationalisation of the binding mode suggested
participation of a putative structural water mol., which was subsequently
confirmed by X-ray crystallog.
RECT (Reactant): SPN (Synthetic preparation)

87769-56-0P

RI: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(arylpyrazolopyridazines as potent inhibitors of glycogen synthase kinase-3)

CAPLUS 87769-56-0

4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-phenyl- (9CI) (CA INDEX NAME)

CORPORATE SOURCE:

REFERENCE COUNT:

THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

136:263177
Preparation of pyridazinones and triazinones exhibiting excellent inhibitory activities against AMPA receptor and/or kainate receptor Magato, Satoshir Kawano, Kokir Ito, Koichir Norimine, Yoshhikor Ueno, Kohshir Hanada, Takahisar Amino, Hiroyukir Jogo, Makotor Hatakeyama, Shinjir Ueno, Masatakar Groom, Anthony John Rivers, Leanner Smith, Terence Bisai Co., Ltd., Japan PCT Int. Appl., 174 pp. CODEN: PIXXD2 PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: Patent Japanese 1 LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PRIORITY APPLN. INFO.: OTHER SOURCE(S):

L4 ANSWER 6 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 2002:220564 CAPLUS DOCUMENT NUMBER: 136:263177
TITLE: Prenaration

L4 ANSWER 6 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

ANSWER 6 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

The title compds. [I; wherein Al, A2 and A3 are each independently C3-8 cycloalkyl, C3-8 cycloalkyl, a 5- to 14-membered anonarom. heterocyclic group, a C6-14 aromatic carbocyclic group, or a 5- to 14-membered aromatic heterocyclic group, any of which may be substituted Q: 5, or NH; Z is C or N; X1, X2 and X3 are each independently a single bond, optionally substituted C1-6 alkylene, optionally substituted C2-6 alkylene, optionally substituted C2-6 alkylene, optionally substituted C2-6 alkylene, optionally substituted C1-6 alkyl, or alternatively R1 and R2 are each independently hydrogen or optionally substituted C1-6 alkyl, or alternatively R1 and R2 may be united in such a way that CR2-ZR1 forms CC; and R3 is hydrogen, optionally substituted C1-6 alkyl, or C2-6 alkynyl, or alternatively R1 may unite with any atom on the ring A1 or A3 to form together with the atom an optionally substituted C1-6 alkyl, c2-6 alkenyl, or C2-6 alkynyl, or alternatively R3 may unite with any atom on the ring A1 or A3 to form together with the atom an optionally substituted C5-8 carbocycle or an optionally substituted S- to 8-membered heterocycle) or salts thereof, or hydrates of both are prepared These compds. do not inhibit N-methyl-D-sapartic acid (NMDA) receptor but they are excellent inhibitors of e-anino-3-hydroxy-5-methyl-4-isomacolepropionic acid (AMPA) receptor and/or kainic acid receptor. They are useful for the prevention or treatment of acute neurodegenerative diseases, acute cerebral vascular disorders, head injury, spinal cord injury, nerve disorders caused by low oxygen or sugar level, chronic neurodegenerative diseases, Altheimer's disease, Parkinson's diseases, Hinhight and the second particle of the property of the pro

L4 ANSWER 7 OF 24 CAPLUS COPYRIGHT 2005 ACS OB STN ACCESSION NUMBER: 1999:576914 CAPLUS DOCUMENT NUMBER: 131:228727

DOCUMENT NUMBER:

131:228727
Preparation of pyridazine derivatives as interleukin 1B production inhibitors ohkuchi, Masao: Kyotani, Yoshinori; Shigyo, Hiromichi; Yoshizaki, Hideo; Koshi, Tomoyukir Kitamura, Takahiro; Matsuda, Takayukir Oda, Soichi; Habata, Yuriko; Kotaki, Kyoko Kowa Co., Ltd., Japan; et al. PCT Int. Appl., 112 pp. CODEN: PIXXO2
Patent
Japanese INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	TENT				KIN		DATE				LICAT				D	ATE		
							1999	0910			1999-				1	9990	226	
	W:	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,	
		DK,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	ΗU,	ID,	IL,	IN,	IS,	JP,	
											LU,							
											, SG,							
											AZ,							T
	RW:										ZV.							
											NL,		SĔ,	BF,	ΒJ,	CF,	ÇG,	
											TD,							
CA	2321	254			AA		1999	0910		CA 1	1999-	2321	254		1	9990	226	
										AU :	1999-	2641	4		1	9990	226	
	7394																	
EP											1999-							
	R:			CH,	DΕ,	DK,	ES,	FR,	GB,	GR,	, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
		IE,																
	5061				λ			1130			1999-					9990		
	2221							0120			2000-					9990		
	6403										-000					0000		
			53							NO 2	2000-	4353			2	0000		
	1035				A1		2004	0820			2001-							
PRIORIT'	Y APP	LN.	INFO	.:							1998-							
										WO :	1999-	JP92	5		W 1	9990	226	

AB The title compds. I (R1 represents lower alkoxy, lower alkylthic or <10/19/2005> Habte

ANSVER 6 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
4-Pyridarinecarboxamide, 2,3-dihydro-N-[2-(4-morpholinyl) ethyl]-3-oxo-6-phenyl- (9Cl) (CA INDEX NAME)

404933-59-1 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-N-[2-(4-morpholiny1)ethy1]-3-oxo-6-phenyl-, monohydrochloride [9CI] (CA INDEX NAME)

• HC1

REFERENCE COUNT:

THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued) halogenos R2 represents H, lower alkoxy, lower alkylthio or halogenos R3 represents OH, CR, halogeno, lower cycloalkyl, lower alkyl or lower alkenyl optionally substituted by an optionally substituted carbamoyl. R4 represents COOH, lower alkoxycorboxyl, optionally substituted amino or optionally substituted are alkoxycorboxyl, optionally substituted amino or optionally substituted ureidos and the dotted line means a single bond or a double bond between the carbon atoms at the 4- and 5-positional are prepat. I are useful as preventives/remedies for immunol. diseases, infismmatory diseases, isohemic diseases, etc. In an in vitro test using cells, 2-cyclopropylmethyl-6-4-methoxyphenyl-4-methylcarbamoyl-2H-pyridazin-3-one showed ICSO of 0.038 µM against lipopolysaccharide-induced interleukin 1 B prodn.
243862-95-59
RL: RCT (Reactant); SFN (Synthetic preparation); PREF (Preparation); RACT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)
(preparation of pyridazine derivs. as interleukin 16 production

inhibitors) RN 243862-95-5 CAPLUS

4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-methoxyphenyl)-N-methyl-3-oxo-(9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 8 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1999:325927 CAPLUS DOCUMENT NUMBER: 130:338106

DOCUMENT NUMBER: TITLE: 130:338106
Preparation of pyrazole derivatives as adenosine Al and A2 antagonists
Akahane, Atsushir Kuroda, Satorus Itani, Hiromichi Pujisawa Pharmaceutical Co., Ltd., Japan PCT Int. Appl., 32 pp.
CODEN: PIXXD2
Patent

INVENTOR (5): PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE WO 9924424 A1 19990520 WO 1998-JF4892 19981028
W: CA, CN, JP, KR, US
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
PT, SE
PRIORITY APPLN. INFO:
OTHER SOURCE(S):
MARPAT 130.338106

MAPPAT 130.338106

MARPAT 130:338106

OTHER SOURCE(S):

The title compds. I [R1 and R2 may be the same or different and each represents optionally substituted aryln R3 represents hydrogen, lower alkyl, or optionally substituted ar(lower) alkyl and R4 represents Q1 (wherein R5 represents optionally substituted ar(lower) alkyl or lower alkanoyl (lower) alkyl), etc.], useful as adenosine A1 and A2 antagonists (no data), are prepared I may serve as preventives and/or remedies for ischemic heart diseases such as angina pectoris, peripheral vascular diseases such as claudication, cerebral ischemia, migraine, diabetes, melancholia, Parkinson's disease, etc. (no data). For example, 3,5-diphenyl-4-[2-(3-methoxybenzyl)-3-oxo-2,3-dihydropyridazin-6-y1]pyrazole was prepared 224573-04-0P
RL: BAC [Biological activity or effector, except adverse); BSU [Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREF (Preparation) USES (Uses) (preparation of pyrazole derivs, as adenosine A1 and A2 antagonists) 224573-04-0 CAPLUS
4-Pyridazinecarboxylic acid, 2,3-dihydro-6-[1-methyl-3,5-diphenyl-lH-pyrazol-4-yl)-3-oxo-, hydrazide (9C1) (CA INDEX NAME)

L4 ANSWER 9 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 1997:558055 CAPLUS DOCUMENT NUMBER: 127:262667 TITLE: Preparation of Title:

127:262667
Preparation of pyrazolo[1,5-a]pyridine derivatives as adenosine antagonists and their pharmaceutical uses Kuroda, Satoshir Itani, Hiromichir Akabane, Atsushi Fujisawa Pharmaceutical Co., Ltd., Japan Jon. Kokai Tokkyo Koho, 17 pp.
CODEN: JXXXAF
Parant

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent

Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DATE PATENT NO. KIND DATE APPLICATION NO. JP 09216883
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
GI A2 JP 1996-24146 JP 1996-24146 19970819 MARPAT 127:262667

The derivs. I [R1 = aryl; R2 = oxodihydropyridazinyl Q [R3 = H, lower alkyl, acyl-lower alkyl, acyl-lower alkyl, acyl-lower alkyl, acyl-lower alkyl, (un) substituted heterocyclyl, (un) substituted lower aralkyl; R4 = H, acyl, cyano, heterocyclyl, lower hydroxyalkyl, (unprotected) anion; R5 = H, lower alkyl; R4 and/or R5 = substituent], pyridazinyl Q1 [R6 = halo, lower alkyl; R4 and/or R5 = substituent], pyridazinyl Q1 [R6 = halo, lower alkyl; R6 = halo, clower alkoxy, (un) substituted anylamino; R7 = acyl, lower hydroxyalkyl]] or their pharmaceutically acceptable salts are claimed. Also claimed are pharmaceuticals containing 1 or their salts and carriers. I show cognition-enhancing, snalgesic, antidepressant, vasodilating, diuretic, cardiotonic, renal circulation-increasing, lipolysis-promoting, antisthmatic, insulin secretion-promoting, platelet aggregation-inhibition; effects, etc., and are especially useful as cardiac infarction inhibitors, antihypertensives, renal failure inhibitors, and diuretics.
3-Propionyl-2-phenylpyrazolol(1,5-alpyridine With (ECO) 20, was successively treated with CO(COZEC)2 at 100 for 65 h then with HZNNHZ.H2O at 125 for 8 h to give 0.42 g 3-(4-(2-isopropylidene)ydrazino) carbony 1-5-methyl-3-oxo-2,3-dihydropyridazin-6-yl]-2-phenylpyrazolol(1,5-alpyridine)

l-5-methyl-3-oxo-4, 3-oinydropyriusein-0-3, 2 p-0-3, 2 p-

their pharmaceutical uses) 195826-98-3 CAPLUS

<10/19/2005>

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ANSWER 8 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
4-Pyridazinecarboxylic acid, 2,3-dihydro-3-oxo-6-(2-phenylpyrazolo[1,5-a)pyridin-3-yl)-, hydrazide (9CI) (CA INDEX NAME)

195827-00-0 CAPLUS 4-Pyridazinecarboxylic acid, 2,3-dihydro-3-oxo-6-(2-phenylpyrazolo[1,5-a]pyridin-3-yl)-, 2-{{1,1-dimethylethoxy}carbonyl}hydrazide (9CI) (CA INDEX NAME)

195827-01-1 CAPLUS 4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(2-phenylpyrazolo[1,5-a]pyridin-3-yl)-N-(3,4,5-trimethoxyphenyl)- (9CI) (CA INDEX NAME)

195827-02-2 CAPLUS Glycine, N-{[2,3-dihydro-3-oxo-6-(2-phenylpyrazolo[1,5-a]pyridin-3-yl}-4-pyridazinyl]carbonyl]-, ethyl ester (9CI) (CA INDEX NAME)

ANSWER 9 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

195827-03-3 CAPLUS
Glycine, N-[[2,3-dihydro-3-oxo-6-(2-phenylpyrazolo[1,5-a]pyridin-3-yl]-4pyridazinyl]carbonyl]- (9CI) (CA INDEX NAME)

195827-04-4 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(2-phenylpyrazolo[1,5-a)pyridin-3-yl)- (9CI) (CA INDEX NAME)

195827-32-8 CAPLUS Carbamic acid, [2,3-dihydro-3-oxo-6-(2-phenylpyrazolo[1,5-a]pyridin-3-yl)-4-pyridazinyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

L4 ANSWER 10 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 1993:625966 CAPLUS 119:225966

TITLE:

INVENTOR(S):

Preparative and biological activity of aryl substituted nitrogen containing heterocycles Linz, Guenter, Fieper, Helmut, Himmelbach, Frank, Austel, Volkhard, Hueller, Thomas, Weisenberger, Johannes, Seewaldt-Backer, Elke Thomae, Dr. Karl, G.m.b.H., Germany Eur. Pat. Appl., 47 pp. CODEN: EFXXDW Patent German 1

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 537696	A1	19930421	EP 1992-117507	19921014
R: AT, BE, CH,	DE, DK	, ES, FR,	GB, GR, IE, IT, LI, LU,	
DE 4134467	A1	19930422	DE 1991-4134467	19911018
US 5418233	A	19950523	US 1992-961135	19921014
CA 2080748	AA	19930419	CA 1992-2080748	19921016
NO 9204027	A	19930419	NO 1992-4027	19921016
AU 9227062	A1	19930422	AU 1992-27062	19921016
AU 662930	B2	19950921		
HU 62272	A2	19930428	HU 1992-3264	19921016
JP 05221992	A2	19930831	JP 1992-277578	19921016
ZA 9207994	A	19940418	ZA 1992-7994	19921016
US 5563268	A	19961008	US 1995-375084	19950119
PRIORITY APPLN. INFO.:			DE 1991-4134467	A 19911018
			US 1992-961135	A3 19921014

PRIORITY APPLM. IMPO.:

DE 1991-4134467 A 1991018

OTHER SOURCE(S):

MARPAT 119:225966

AB The preparation of title compds. with fibrinogen-binding, thromboxane, and blood platelet aggregation inhibitor activity is claimed. Thus, reaction of 6-(4-amidinopheny)1-4-[4-(methoxycarbony)1buy]1 aminocarbony]1-2-methy1-(2H)-pyridazin-3-one (preparation given) with LiOH.H2O in a mixture

THF-H2O gave 91.1 6-(4-amidinophenyl)-4-[(4-carboxybutyl)aminocarbonyl]-2-methyl-(2H)-pyridazin-3-one. Similarly, a number of pyridazinone and pyrimidine derivs. were prepared and their biol. activity is described. 150594-47-19

IT 150594-47-1P
RL: RCT (Reactant), SPN (Synthetic preparation), PREP (Preparation), RACT (Reactant or reagent)
Inhibitor)
RN 150594-47-1 CAPLUS
CN Pentanoic acid, 5-11/6-1/

bitor)
150594-47-1 CAPLUS
Pentanoic acid, 5-[[[6-(4-cyanophenyl)-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]- [9CI] (CA INDEX NAME)

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L4 ANSWER 9 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

150594-75-5P 150594-91-5P 150595-00-9P
150595-14-5P 150595-38-3P
RL: SPM (Synthetic preparation), PREP (Preparation)
(preparation and thromboxane formation inhibiting activity of)
150594-75-5 CAPLUS
Pentanoic acid, 5-[[[6-[4-,(aminoiminomethyl)phenyl]-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]- (9CI) (CA INDEX NAME) ΙT

150594-91-5 CAPLUS Cyclohexanecarboxylic acid, 4-[[[6-[4-(aminoiminomethyl)phenyl]-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]-, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

150595-00-9 CAPLUS Pentancic acid, 5-[[[6-[4-(aminoiminomethyl)phenyl)-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]-, methyl ester (9CI) (CA INDEX NAME)

150595-14-5 CAPLUS Cyclohexanecarboxylic acid, 4-{[[6-{4-{aminoiminomethyl}phenyl]-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]-, methyl ester, hydrochloride, trans- (9CI) (CA INDEX NAME)

L4 ANSWER 10 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN Relative stereochemistry. (Continued)

150595-38-3 CAPLUS Cyclohexanecarboxylic acid, 4-[{6-[4-(aminoiminomethyl)phenyl}-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]-, methyl ester, trans- (9CI) (CA INDEX NAME)

ANSWER 11 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
Thus, 4-(5-cyano-2-pyridyl)phenol (prepn. given) was condensed with
(35,55)-3-[(tert-butyloxycarbonyl)methyl)-5-[(methanesulfonyloxy)methyl]-2pyrrolidinone and the product converted in 2 steps to title compd.
(35,55)-1 which had EO50 of 0.06 µM against collagen-induced platelet
aggregation in vitro.
149354-60-9P 149354-62-1P 149354-79-0P
149354-61-9P 149355-3-PP 149377-23-1P
RL: BRC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SFN (Synthetic preparation); BIOL (Biological
study); PREP (Preparation)
(preparation of, as cell aggregation inhibitor)
149354-60-9 CAPLUS
3-Pyrrolidineacetic acid, 5-[[[6-[4-(aminoiminomethyl)phenyl]-2,3-dihydro3-cxo-4-pyridazinyl]carbonyl]amino|methyl]-2-oxo-, (35-trans)- (9CI) (CA
INDEX NAME)

Absolute stereochemistry.

149354-62-1 CAPLUS
3-Pyrrolidineacetic acid, 5-[[[[6-[4-(aminoiminomethyl)phenyl]-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]methyl]-2-oxo-1-(3-phenylpropyl)-,
[35-trans]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

149354-79-0 CAPLUS
3-Pyrrolidineacetic acid, 5-[[[[6-[4-(aminoiminomethyl)phenyl]-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]methyl]-2-oxo-1-(3-phenylpropyl)-,
methyl ester, monohydrochloride, (35-trans)- (9CI) (CA INDEX NAME)

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L4 ANSWER 11 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1993:517098 CAPLUS DOCUMENT NUMBER: 119:117098

TITLE:

INVENTOR(S):

119:117098 CAPLUS
119:117098 Preparation of 2-pyrrolidinone-3-acetates and analogs as cell aggregation inhibitors
Austel, Volkhard; Eisert, Wolfgang; Himmelsbach,
Frank; Linz, Guenter; Hueller, Thomas; Pieper, Helmut;
Weisenberger, Johannes
Thomae, Dr. Karl, G.m.b.H., Germany
Eur. Pat. Appl., 73 pp.
CODEN: EPXKDW
Patent
German

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.		DATE	APPLICATION NO.	
EP 528369	A2	19930224	EP 1992-113877	19920814
EP 528369	A3	19930421		
EP 528369	B1	19991124		
		, ES, FR,	GB, GR, IE, IT, LI, LU,	NL, PT, SE
DE 4127404	A1	19930225	DE 1991-4127404	19910819
AT 186906	E	19991215	AT 1992-113877	19920814
CA 2076311	AA	19930220	CA 1992-2076311	19920818
NO 9203235	A	19930222	NO 1992-3235	19920818
AU 9221119	A1	19930225	AU 1992-21119	19920818
AU 654372	B2	19941103		
JP 06025227	A2	19940201	JP 1992-219149	19920818
ZA 9206205	A	19940218	ZA 1992-6205	19920818
IL 102847	A1	19961114	IL 1992-102847	19920818
US 5455348	A	19951003	US 1993-173603	19931223
PRIORITY APPLN. INFO.:			DE 1991-4127404	A 19910819
			US 1992-929870	B1 19920814
OTHER SOURCE(S):	MARPAT	119:11709		

$$\begin{array}{c|c} \text{H}_2\text{N} & \text{CH}_2\text{CO}_2\text{H} \\ \hline \end{array}$$

EYAXIXZX3X4X5B [A = (substituted) bivalent (oxo)alkyleneimino; B = NH2, C(:NH)NH2, NHC(:NH)NH2, etc.; E = CO2H, alkoxycarbonyl, etc.; X1 = bond, alkylene; X2 = bond, O, NH, SO2NH, etc.; X3, X5 = (heterojcycloalkylene, (heterojarylene, etc.; X4 = bond, O, CH2, CO, NH, etc.; X3XXF = phenylene, (CH2)3-5, etc.; Y = alkylene, NHCH2, OCH2, etc.] were prepared

L4 ANSWER 11 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN Absolute stereochemistry. (Continued)

● HC1

149355-41-9 CAPLUS
3-Pyrrolidineacetic acid, 5-[[[[6-(4-cyanopheny1)-2,3-dihydro-3-oxo-4-pyridaziny1]carbony1]amino]methyl]-2-oxo-, methyl ester, (35-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

149355-53-3 CAPLUS
3-Pyrrolidineacetic acid, 5-[{[[6-(4-cyanophenyl)-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]methyl]-2-oxo-1-(3-phenylpropyl)-, methyl ester, (3S-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

10/715,556

Page 17

L4 ANSWER 11 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

149377-23-1 CAPLUS
3-Pyrrolidineacetic acid, 5-{{[[6-[4-{aminoiminomethyl]phenyl]-2,3-dihydro-3-oxo-4-pyridazinyl]carbonyl]amino]methyl]-2-oxo-, methyl ester, monohydrochloride, (3S-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 2-A • HC1

ANSWER 12 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (9CI) (CA INDEX NAME) (Continued)

L4 ANSWER 12 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STR ACCESSION NUMBER: 1991:206976 CAPLUS DOCUMENT NUMBER: 114:206976 114:206976
Synthesis of aza analogs of amrinone
Singh, Baldev, Lesher, George Y.
Dep. Hed. Chem., Sterling Res. Group, Rensselaer, NY,
12144, USA.
Heterocycles (1990), 31(12), 2163-72
CODDM: HTCYAM; ISSN: 0305-5414 TITLE: AUTHOR(S): CORPORATE SOURCE:

SOURCE: Journal English CASREACT 114:206976 DOCUMENT TYPE:

OTHER SOURCE(S):

The aldol condensation product I of 4-acetylpyridine and CO(COZEt)2 was converted to pyridazinecarboxylic acid hydrazide II (R = CONHMHZ)(III). Curtius reaction of III gave aminopyridazinone II (R = NHZ). The condensation of (4-pyridyl)glyoxal with aminomalonamide HZNCH(COMHZ)2 yielded pyrazinecarboxamide IV (R1 = CONHZ) which was transformed to aminopyrazinone IV (R1 = NHZ) by the Hofmann reaction. Curtius reaction of 1,2,4-triazinone-5-carboxylic acid V (R2 = COZH) gave aminotriazinone V (R2 = NHZ). Demethylation of methoxypyrimidine VI (R3 = He) gave pyrimidinol VI (R3 = H).
80843-46-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and Curtius reaction of)
80843-46-5 CAPLUS
4-Pyridazinecarboxylic acid, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-, hydrazide

L4 ANSWER 13 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1990:91460 CAPLUS
112:91460 TITLE: 112:91460

AUTHOR(S): Pyridazines. Part 43. Pyridazine analogs of biologically active compounds. Part 5: Novorl potential cardiotonics of the amrinone type Haider, N., Hainisch, G.; Offenberger, Sigrid inst. Pharm. Chem., Univ. Vienna, Vienna, A-1090, Austria

SOURCE: Pharmazie (1989), 44(9), 598-601 CODEN: PHARAT; ISSN: 0031-7144

DOCUMENT TYPE: LANGUAGE: English

DOCUMENT TYPE: LANGUAGE: GI

Preparation of a series of novel pyridazine derivs. structurally related to bipyridine cardiotonics, starting from 4-methylpyridazine or 4-methylpyridazine, resp., is described. As observed with I, II and III, an enhancement of in vitro cardiotonic activity was associated with the replacement of one or both pyridine subunit(s) in amrinone or milrinone by a 1,2-diazine system.

125375-18-0P

RL: RCT (Reactant); SFN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and Hofmann degradation of)

125375-18-0 CAPLUS [3,4"-Bipyridazine)-5-carboxamide, 1,6-dihydro-6-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 14 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
111:97259 CAPLUS
111:97259
111:97259
111:97259
111:97259
111:97259
Preparation of phenylpyridazinone derivatives as cardictonics and antihypertensives
Sircar, 11as Bristol, James A.
Varner-Lambert Co., USA
U.S., 19 pp. Cont.-in-part of U.S. Ser. No. 407,973.
CODEN: USXXAM
DOCUMENT TYPE:
FAMILY ACC. NUM. COUNT:
FAMILY ACC. NUM. COUNT:

FAMILY ACC. NUM. COUNT:

FAMILY ACC. NUM. COUNT:

ACCESSION NUMBER:
1199:497259 CAPLUS
1199:497259 CAP

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
				-	
US 4734415	A	19880329	US 1983-477695		19830322
US 4353905	Α	19821012	US 1981-302181		19810917
PRIORITY APPLN. INFO.:			US 1981-302181	A2	19810917
			US 1982-402488	A2	19820727
			US 1982-407973	A2	19820813

OTHER SOURCE(S): CASREACT 111:97259; MARPAT 111:97259

$$\sum_{Y} \sum_{N=N/2}^{R^3} x^4$$

The title compds. [I; dotted line represents single or double bond; X = 0, S; R2 = H, lower alkyl; R3 = H, lower alkyl; when dotted line represents a single bond, R3 = dilower alkyl; R4 = H, lower alkyl; or when dotted line represents a double bond, R4 = H, lower alkylamino, cyano, OM, CHZOH, CONNSR6, etc.; R3R4 = atoms to complete a carbocycle of 3-6 atoms; R5, R6 = H, alkyl Y = H, halo, lower alkyl, alkoxy etc.; A = R12; R1 = N-attached, (un)substituted, 5- or 6-membered heterocyclyl, optionally containing other hetero atoms; Z = bond, (CHZ)nO in the 4-position; n = 2-5] and their pharmaceutically acceptable salts, useful as cardiotonics and antihypertensives, were prepared 97150-66-R9 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and reaction of, in preparation of cardiotonic and hypertensive)

antihypertensive)
RN 97150-66-8 CAPLUS
CN 4-Pyridazinecarboxylic acid, 2,3-dihydro-6-[4-(1H-imidazol-1-yl)phenyl]-3oxo-, hydrazide (9CI) (CA INDEX NAME)

L4 ANSWER 15 OF 24 CAPLUS COPYRIGHT 2005 ACS OR STN ACCESSION NUMBER: 1989:114776 CAPLUS DOCUMENT NUMBER: 110:114776

110:114776
3-Aminopyridazine derivatives with atypical antidepressant, serotonergic and dopaminergic

activities
Wermuth, Camille Georges; Schlewer, Gilbert; AUTHOR(S):

Wermith, Camille Georges; Schlewer, Gilbert; Bourquignon, Jean Jacques; Maghioros, Georges; Bouchet, Marie Jeanne; Moire, Claudine; Kan, Jean Paul; Worms, Paul; Biziere, Kathleen Dap. Pharmacochim. Mol., Univ. Louis Pasteur, Strasbourg, 67084, Fr. Journal of Medicinal Chemistry (1989), 32(3), 528-37 CODEN: JMCMAR; ISSN: 0022-2623

CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE: LANGUAGE:

Journal English CASREACT 110:114776 OTHER SOURCE (S):

Forty-seven substituted analogs of minaprine, e.g., I, were synthesized and tested for their potential antidepressant, serotonergic, and dopaminergic activities. The structure-activity relationships show that dopaminergic and serotonergic activities can be dissociated Serotonergic activity appears to be correlated mainly with the substituent in the 4-position of the pyridazine ring whereas the dopaminergic activity appears to be dependent on the presence, or in the formation, of a para-hydroxylated aryl ring in the 6-position of the pyridazine ring. 8759-56-OP

L4 ANSWER 14 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

97150-66-8
RL: RCT (Reactant): RACT (Reactant or reagent)
(reaction of, in preparation of pyridazinone cardiotonic and
antihypertensive)
97150-66-8 CAPLUS
4-Pyridazinecarboxylic acid, 2,3-dihydro-6-[4-(1H-imidazol-1-yl)phenyl]-3-oxo-, hydrazide (9CI) (CA INDEX NAME)

L4 ANSWER 16 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1987:598252 CAPLUS DOCUMENT NUMBER: 107:198252 CAPTUS CAPT

139.1398222
Cardiotonic agents. 7. Inhibition of separated forms of cyclic nucleotide phosphodiesterase from quinea pig cardiac muscle by 4.5-dihydro-6-[4-(IH-imidazol-1-yl)phenyl]-3(2H)-pyridazinones and related compounds. Structure-activity relationships and correlation with in vivo positive inotropic activity
Siccar, Ila, Weishaar, Ronald E., Kobylarz, Dianne, Moos, Walter H., Bristol, James A.
Dep. Chem., Warner-Lambert/Parke-Davis Pharm. Res., Ann Arbor, MI, 48105, USA
Journal of Medicinal Chemistry (1987), 30(11), 1955-62
CODEN: JMCMAR; ISSN: 0022-2623
Journal
English
CASREACT 107:198252

AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE: LANGUAGE:

OTHER SOURCE(S):

$$\begin{bmatrix} N \\ M \end{bmatrix} \longrightarrow \begin{bmatrix} N \\ N \\ M \end{bmatrix} \longrightarrow$$

Imidazolylphenylpyrazolinone I was prepared from benzonitrile II. The structure-activity relationships of a series of 4,5-dihydro-6-[4-(1H-imidazol-1-yl)phenyl]-3(2H)-pyridazinones, e.g., III (R = H, Me, CH2Th, CH2CH2OH, CH2CH2OAC, RI = H, Me, MH2, CONNE; R2 = H, He, Et R3 = H, He, SH, SNe, SOMe, Et), I and related compds. were investigated for the in vivo inhibition of different forms of cyclic nucleotide phosphodiesterase (PDE) isolated from guines pig ventricular muscle. With few exceptions, these 4,5-dihydropyridazinones were potent inhibitors of cardiac type III phosphodiesterase, which is a low Km, CAMP specific form of the enzyme. The inhibitory effects on cardiac type I and type II phosphodiesterase, both of which hydrolyze CAMP as well as cyclic GH, were minimal. The most selective PDE III inhibitor was CI-930 III (R = R1 = R3 = H, R2 = Me) (IV), the 5-Me analog of imazodan III (R = R3 = H) with an EDS of 0.6 µH. The most potent inhibitor of PDE III was the 4,5,6,7-tetrahydrobenzimidazole analog of IV, with an EDS of 0.15 µM. The structural features that impert both selectivity for inhibiting type III phosphodiesterase and potency of inhibition and correlations between in vitro PDE inhibitory potency, in vivo pos. inotropic potency, and physicochem. properties are discussed.

87150-66-8 97150-67-9
RL: RCT (Reactant) RACT (Reactant or reagent) (phosphodiesterase inhibitory activity of) 87150-66-8 CAPLUS 4-Pyridazinecarboxylic acid, 2,3-dihydro-6-[4-(1H-imidazol-1-yl)phenyl]-3-oxo-, hydrazide (9CI) (CA INDEX NAME)

10/715,556

Page 19

L4 ANSWER 16 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

97150-67-9 CAPLUS 4-Pyridazinecarboxamid (9CI) (CA INDEX NAME) oxamide, 2,3-dihydro-6-[4-(1H-imidazol-1-yl)phenyl]-3-oxo-

L4 ANSWER 18 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 1985:560462 CAPLUS DOCUMENT NUMBER: 103:160462 CAPTULE: Cardiotopic access to the control of the cont

Cardiotonic agents. 2. Synthesis and structure-activity relationships of 4,5-dihydro-6-[4-(H-imidazol-1-yl)phenyl]-3(2H)-pyridazinones: a new class of positive inotropic agents sircar, Ila; Duell, Bradley L.; Bobowski, George; Bristol, James A.; Evans, Dale B. Dep. Chem., Warner-Lambert/Parke-Davis Pharm. Res., Ann Arbor, M.; 48105, USA
Journal of Medicinal Chemistry (1985), 28(10), 1405-13 CODEN: JMCMAR; ISSN: 0022-2623
Dournal English

AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE:

LANGUAGE: OTHER SOURCE(S): GI English CASREACT 103:160462

A series of 4,5-dihydro-6-[4-(lH-imidazol-1-yl)phenyl]-3(2H)-pyridazinones and related compds. were synthesized and evaluated for pos. inotropic activity. Most members of this series produced dose-related increases in myocardial contractility that were associated with relative minor increases in heart rate and decreases in systemic arterial blood pressure. Introduction of a Me substituent at the 5-position of pyridazinone I (R = H) (II) produced the most potent compound in this series, I (R = He) (III). Compound II is more potent than amrinone whereas compound III is more potent than mininone. The inotropic effects of II and III are not mediated via stimulation of B-adrenergic receptors. Selective inhibition of cardiac phosphodiesterase fraction III represents the principal component of the pos. inotropic action of II and III. 97180-67-99
RIL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SRN (Synthetic preparation); BIOL (Biological study), PREP (Preparation)
(preparation and inotropic activity of)
97150-67-9 CALIUS
4-Pyridazinecarboxamide, 2,3-dihydro-6-[4-(lH-imidazol-1-yl)phenyl]-3-oxo-(9CI) (CA INDEX NAME)

<10/19/2005>

Habte

L4 ANSWER 17 OF 24 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 1987:213880 CAPLUS DOCUMENT NUMBER: 106:213880

TITLE:

106:213880
The reaction of pyridazinones with nucleophiles. An unusual reaction with cyanide
Badger, Edward W., Moos, Walter H.
Dep. Chem., Warner-Lambert/Parke-Davis Pharm. Res.,
Ann Arbor, HI, 48105, USA
Journal of Heterocyclic Chemistry (1986), 23(5),
1515-17
CODEN., NECKEN. 1682. AUTHOR (S): CORPORATE SOURCE: SOURCE:

CODEN: JHTCAD: ISSN: 0022-152X

Journal

LANGUAGE: OTHER SOURCE(5): English CASREACT 106:213880

DOCUMENT TYPE:

Studies on the synthesis of pyridazinone analogs of pyridone cardiotonics are reported. The synthetic scheme involves the reaction of pyridazinones and chloropyridazinones I (R=R,R=H,C) with nucleophiles. Addition occurred twice with cyanide as the nucleophile, thus providing a novel dicyanopyridazinone I (R=R1 = cyano).

ΙŤ

97150-66-6
RL: RCT (Reactant): RACT (Reactant or reagent)
(Curtis rearrangement of)
97150-66-8 CAPLUS
4-Pyridazinecarboxylic acid, 2,3-dihydro-6-[4-(1H-imidazol-1-yl)phenyl]-3-cxo-, hydrazide (9CI) (CA INDEX NAME)

ANSWER 18 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

97150-66-8P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
[preparation, hydrolysis, and inotropic activity of)
97150-66-8 CAPLUS
4-Pyridazinecarboxylic acid, 2,3-dihydro-6-[4-(1H-imidazol-1-y1)phenyl]-3-oxo-, hydrazide (9CI) (CA INDEX NAME) IT

L4 ANSWER 19 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1985:62255 CAPLUS DOCUMENT NUMBER: 102:62255

DOCUMENT NUMBER:

102:62255
Pyridazine derivative having a psychotropic action and medicines containing them
Kan, Jean Paulr Biziere, Kathleen, Wermuth, Camille Georges
Sanofi, Fr.
Fr. Demande, 12 pp.
CODEN: FROXBL
Patent TITLE:

INVENTOR (S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: Patent French

KIND	DATE	APPLICATION NO.	
A1		FR 1983-1366	19830128
Bl	19850607		
λ	19860121	US 1984-571696	19840118
A1	19870303	CA 1984-445482	19840118
A	19840729	DK 1984-259	19840120
В	19910107		
C	19910527		
A	19840829	ZA 1984-500	19840123
A1	19870331	IL 1984-70755	19840123
A1	19840802	AU 1984-23728	19840124
B2	19871015		
A1	19841001	ES 1984-529108	19840124
A1	19840822	EP 1984-400157	19840125
B1	19880127		
, DE, FI	, GB, IT, L	I, LU, NL, SE	
E	19880215	AT 1984-400157	19840125
A	19840729	FI 1984-349	19840127
В	19881130		
С	19890310		
A	19840730	NO 1984-329	19840127
0	19841029	HU 1984-378	19840127
В	19870828		
A5	19841114	DD 1984-259679	19840127
A3	19861130	SU 1984-3697653	19840127
B1	19880430	PL 1984-245932	19840127
B2	19910411	CS 1984-614	19840127
A2	19840814	JP 1984-14185	19840128
A	19861223	US 1985-735580	19850520
Ä	19891208	DK 1989-6215	19891208
R	19910930		
č	19920302		
Ă	19891208	DK 1989-6216	19891208
R	19910930		
ř	19920302		
	13320002	FR 1983-1366	A 19830128
		FR 1983-18433	A 19831118
		IIS 1984-571696	A3 19840118
		ED 1984-400157	A 19840125
CASRE	ACT 102:6225		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
	A B C A A1 B2 A1 BB C A B B C	All 19840803 Bl 19850607 A 19860121 Al 19870303 A 19840729 B 19910107 C 19910527 A 19840829 Al 19840802 Bl 19871015 Al 19840802 Bl 19871015 Al 19840802 Bl 19871015 Al 19840802 Bl 19871015 Al 198408012 C 19890310 A 19840730 C 19890310 A 19840730 D 19841001 Al 19840730 D 19841001 Al 19840730 C 19890310 A 19840730 D 1984030 Bl 19880430 Bl 19880430 Bl 19880430 Bl 19880430 Bl 1980830 C 19920302 A 1986132 A 1986134 A 1986123 A 19991083 B 19910930 C 19920302 A 19891208 B 19910930 C 19920302	Al 19840803 FR 1983-1366 Bl 19850607 A 19860121 US 1984-571696 Al 19870303 CA 1984-445482 A 19840729 DK 1984-259 B 19910107 C 19910527 A 19840829 2A 1984-500 Al 19870331 IL 1984-70755 Al 19840829 AU 1984-23728 B2 19871015 Al 19840802 EP 1984-2728 B1 19840802 EP 1984-400157 B1 19840127 B1 19860127 C, DE, FR, GB, IT, LI, LU, NL, SE E 19820215 AT 1984-400157 A 19840729 FI 1984-349 B 1981130 C 19890310 A 19840730 NO 1984-329 O 19841029 HU 1984-378 B 19870828 AS 19841114 DD 1984-378 B 19870828 AS 19841114 DD 1984-259679 AS 19840730 VL 1984-259679 AS 19840130 SU 1984-3697653 Bl 19800430 PL 1984-245932 B2 19910411 CS 1984-6014 A 19861223 US 1984-3575580 B 19910930 C 19920302 A 19891208 DK 1989-6216 B 19910930 C C 19920302 FR 1983-1366 FR 1983-1366 FR 1983-1366 FR 1983-1366 FR 1983-1366 FR 1983-136433 US 1984-511656

L4 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1985:24642 CAPLUS
DOCUMENT NUMBER: 102:24642
ITITLE: 102:24642
INVENTOR(S): Biziere, Kathleen; Kan, Jean Paul; Wermuth, Camille Georges
PATENT ASSIGNEE(S): Sanofi, Fr.
SOURCE: EUR. Pat. Appl., 29 pp.
COUNT EPXKDW
DOCUMENT TYPE: Patent

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: French 2

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
EP 116494	A1 19840822	EP 1984-400157	19840125
EP 116494	B1 19880127		
R: AT, BE, CH,	DE, FR, GB, IT, LI,	LU, NL, SE	
FR 2540115	A1 19840803	FR 1983-1366	19830128
FR 2540115	B1 19850607		
FR 2555178	A1 19850524	FR 1983-18433	19831118
FR 2555178	B1 19860418		
AT 32220	E 19880215	AT 1984-400157	19840125
PRIORITY APPLN. INFO.:		FR 1983-1366 A	19830128
		FR 1983-18433 A	19831118
		EP 1984-400157 A	19840125
OTHER SOURCE(S):	CASREACT 102:24642		

3-Amino-4-pyridazinecarbonitriles I [one of R and Rl is H or alkyl, and the other is H, alkyl, cycloalkyl, Ph or substituted Ph, naphthyl, thienyl, 3-indolyl, Z = CHZCHZ, CHZCHMe, (CHZ)3, RZ = H and R3 = H, CHZCHZO, NO NNZR3 = 4-morpholinyl, 3-mox-4-morpholinyl), which were prepared, showed psychotropic activity. 3-Chloro-6-phenyl-4-pyridazinecarbonitrile was heated with 4-(2-aminoethyl)morpholine in BuOH to give 1 (R = Ph, R1 = H, Z = CHZCHZ, NRZR3 = 4-morpholinyl). RACT (Reactant), SPN (Synthetic preparation), PREP (Preparation), RACT (Resctant or reagent) (Preparation and reaction of, with phosphoryl chloride) 87769-56-0 CAPLUS (4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-phenyl- (9CI) (CA INDEX NAME) AB

ANSWER 19 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

3-[2-(4-Morpholinyl) ethylamino]-6-phenyl-4-pyridazinecarbonitrile dihydrochloride [I] was prepared, and it showed antidepressant activity. The cyclocondensation of PhCOCH2CH(CO2Et)2 with N2H4 gave pyridazinone derivative II, which was brominated and dehydrobrominated to give ester III

- ORt), the latter was converted to anide III (R = NH2). The amide was treated with POC13 to give 3-chloro-6-phenyl-4-pyrazinecarbonitrile, and the product was treated with 4-(2-aminosthyl)morpholine and HCl to give I. 8769-86-0P (R IT

87769-56-0P
RL: RCT (Reactant), SPN (Synthetic preparation), PREP (Preparation), RACT (Reactant or reagent)
(preparation and dehydration of, by phosphoryl chloride)
87769-56-0 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-phenyl- (9CI) (CA INDEX NAME)

ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

94011-51-5 94011-52-6 94011-53-7
94011-54-8 94011-55-9 94011-56-0
94011-57-1 94011-58-2 94011-59-3
94011-60-6 94011-61-7 94011-62-8
94011-63-9
RL: RCT (Reactant), RACT (Reactant or reagent)
(reaction of, with phosphoryl chloride)
94011-51-5 CAPLUS
4-Pyridazinecarboxamide, 6-(4-chlorophenyl)-2,3-dihydro-3-oxo- (9CI) (CA
INDEX NAME)

94011-52-6 CAPLUS 4-Pyridazinecarboxamide, 6-cyclohexyl-2,3-dihydro-3-oxo- (9CI) (CA INDEX NAME)

94011-53-7 CAPLUS 4-Pyridazinecarboxamide, 6-(2,4-dichlorophenyl)-2,3-dihydro-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 94011-54-8 CAPLUS
CN 4-Pyridazinecarboxamide, 6-(2-chloropheny1)-2,3-dihydro-3-oxo- (9CI) (CA INDEX NAME)

RN 94011-55-9 CAPLUS CN 4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-(2-thienyl)- (9C1) (CA INDEX NAME)

RN 94011-56-0 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-methoxyphenyl)-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 94011-60-6 CAPLUS CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-methylphenyl)-3-oxo- (9CI) (CA INDEX NAME)

RN 94011-61-7 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-[4-(trifluoromethyl)phenyl](SCI) (CA INDEX NAME)

RN 94011-62-8 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-[3-(trifluoromethyl)phenyl](SCI) (CA INDEX NAME)

RN 94011-63-9 CAPLUS
CN 4-Pyridazinecarboxamide, 6-(4-cyanophenyl)-2,3-dihydro-3-oxo- (9CI) (CA HNEX NAME)

L4 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

RN 94011-57-1 CAPLUS
CN 4-Pyridazinecarboxamide, 6-(3,4-dichlorophenyl)-2,3-dihydro-3-oxo-(9CI)
(CA INDEX NAME)

RN 94011-58-2 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-{2-naphthalenyl}-3-oxo- (9CI) (CA INDEX NAME)

RN 94011-59-3 CAPLUS
CN 4-Pyridazinecarboxamide, 2,3-dihydro-6-(4-nitrophenyl)-3-oxo- (9CI) (CA INDEX NAME)

L4 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

L4 ANSWER 21 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1984:630453 CAPLUS
DOCUMENT NUMBER: 101:230453
ITILE: Novel anniantion of 6-aryl-3(2H)-pyridazinones with hydrazine
SIngh, Baldev
Sterling-Winthrop Res. Inst., Rensselser, NY, 12144, USA
SOURCE: Heterocycles (1984), 22(8), 1801-4
CODEN: HTCYAM: ISSN: 0385-5414
JOURNAL LANGUAGE: English
OTHER SOURCE(S): CASREACT 101:230453

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

Aminopyridazinones I (R = H, He; Rl = 4-pyridyl, 4-H2NCGH4, 4-HOCGH4) were prepared from II (R2 = 4-pyridyl, 4-AcKHCGH4, 4-HOCGH4). II (R = H, R2 = 4-pyridyl) was heated with N2H4 to give I (R = H, Rl = 4-pyridyl).

80843-46-5

AL: RCT (Reactant); RACT (Reactant or reagent) (attempted rearrangement of, with hydrazine)

80843-46-5

CAPUIS

4-Pyridazinearboxylic acid, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-, hydrazide (9CI) (CA INDEX NAME)

L4 ANSWER 23 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1983:594988 CAPLUS
DOCUMENT NUMBER: 99:194988
Substituted 6-phenyl-3(ZH)-pyridazinones useful as cardiotonic agents
SIrcar, 11a
Varner-Lambert Co., USA
U.S., 6 pp.
COMENT TAND.

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 1981-263643 US 1981-325719 19810514 19830913 US 4404203 US 4397854 19830809 PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI US 1981-263643 CASREACT 99:194988

The cardiotonic title compds. I [R = H, alkyl, PhCH2, Ph; Rl = H, R2 = CF3, PhCH2, cyano, CO2H, CONR52 (R5 = H, alkyl), CH2NR52, CH2CH, NR52; R2 = H, Rl = CF3, cyano, CONR52, CH2NR52, NR52; R3 = H, halo, alkyl, slkony, H0, PhO, sulfonanido; dotted line represents single or double bond) were prepared This SP PhCCH2CH2CO2H was cyclized with H2NH12.H2O in Itol to give 75.6 g 6-phenyl-4,5-dihydro-3(2H)-pyridazinone, which was dehydrogenated by treatment with Br to give 60 g 6-phenyl-3(2H)-5-24,5-dihydro-3(2H)-5-3(2H)

ΙŢ

L4 ANSWER 22 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 1984:174757 CAPLUS
100:174757
AUTHOR(S): Sircar, 1la separal procedure
Sircar, 1la Dep. Chem., warner-Lambert/Parke-Davis Pharm. Res., Ann Arbor, MI, 48105, USA
JOURNAL OF COMENT TYPE: 1000-11473-6
CODEN: JHTCAD; ISSN: 0022-152X
JOURNAL OF COMENT TYPE: 1984:17475-6
CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE:

LANGUAGE: OTHER SOURCE(S): English CASREACT 100:174757

3,4-Dichloro-6-phenylpyridazine (1) was prepared by treating
2-benzyl-4,5-dihydro-6-phenyl-3(2H)-pyridazinone with PCIS-POCI3. I was
aminated to give II [R = NMe2, NH(CH2)3NMe2, NHBU, 4-methylpiperizino,
morpholino, thiomorpholino; Rl = Cl] which were hydrolyzed with acid to II
(Rl = OH).
87769-56-0P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation and Hofmann degradation of)
87769-56-0 CAPLUS
4-Pyridazinecarboxamide, 2,3-dihydro-3-oxo-6-phenyl- (9CI) (CA INDEX
NAME)

L4 ANSWER 23 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	KIND	DATE	APPLICATION NO.		DATE
FR 2481284	A1	19811030	FR 1981-8251		19810424
US 4304776	A	19811208	US 1980-144697		19800428
US 4305943	A	19811215	US 1980-144563		19800428
US 4338446	A	19820706	US 1981-238483		1981022
US 4346221	λ	19820824	US 1981-239566		1981030
AU 8169724	A1	19811105	AU 1981-69724		1981042
GB 2075500	λ	19811118	GB 1981-12638		1981042
GB 2075500	B2	19840606			
ZA 8102652	A	19820526	ZA 1981-2652		1981042
BE 888566	A1	19811027	BE 1981-10209		1981042
DK 8101866	A	19811029	DK 1981-1866		1981042
FI 8101304	A	19811029	FI 1981-1304		1981042
NO 8101420	A	19811029	NO 1981-1420		1981042
SE 8102660	A	19811029	SE 1981-2660		
ES 501665	A1	19830101	ES 1981-501665		1981042
CA 1166253	A1	19840424	CA 1981-376309		1981042
CA 1166254	A1	19840424	CA 1981-376317		1981042
NL 8102077	λ	19811116	NL 1981-2077		1981042
JP 56167684	A2	19811223	JP 1981-65103		1981042
DE 3116861	A1	19820114	DE 1981-3116861		1981042
RIORITY APPLN. INFO.:			US 1980-144563	A	19800429
			US 1980-144697	A	1980042
THER SOURCE(S):	CASRE	ACT 96:85571			

L4 ANSWER 24 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

Cardiotonic (no data), pyridylpyridazinones I (R = H, alkyl, hydroxyalkyl; R1 = NHZ, CONHZ, COZH, CONHNHZ, alkoxycarbonyl) were prepared Thus 4-acetylpyridine was treated with OC(COZE1) 2 to give II which was cyclized with NZH4 and dehydrated to give III (R = COZEL). The ester was converted to the hydrazide and then the azide which was subjected to Curtius rearrangement, hydrolysis, and decarboxylation to give III (R1 = NHZ).

80843-46-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and reaction of, with nitrite)
80843-46-5 CAPLUS
4-Pyridazinecarboxylic acid, 2,3-dihydro-3-oxo-6-(4-pyridinyl)-, hydrazide (9CI) (CA INDEX NAME)